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(10) (CA) CANADIAN PATENT (12)

- (5A) Process for Activating Collulose Pulp with NO and/or NO₂ and Oxygen in the Presence of Nitric Acid and Then Delignifying Bleaching the Activated Cellulose Pulp
- (72) Samuelson, Kans O., Sweden
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A PROCESS FOR ACTIVATING CELLULOSE PULL WITH NO AND/OR MO, AND OXYGEN IN THE PRESENCE OF NITTIC ACID AND THEN DELIGNIFYING BLEACHING THE ACTIVATED CELLULOSE PULP

ABSURACT OF THE DISCLOSURE

A process is provided for activating cellulose pulp using NO and/or NO, plus expect gas in the presence of nitric acid, edded in an amount within the range from about 0.1 to about 1.0 gmole per kg of water accompanying the cellulose gulp at a temperature within the range from about 40 to about 130°C for a time at an activating temperature of 40 to 50°C of from 15 to 180 minutes, at from 50 to 30°C of from 5 to 120 minutes, and at higher temperatures of from 1 to 10 minutes, tollowed by washing, and delignifying bloaching in an alkaline medium with or without expense gas and/or peroxide.

Having regard to the foregoing disclosure, the following is claimed as the inventive and petomable embodiments thereof:

- delignifying bloaching the activated pulp, which comprises treating the cellulose pulp in an activating stage with altrogen oxide selected from the group consisting of altrogen dioxide, mixagen monoxide, pulymers thereof, and mixtures thereof, and with an oxygen-containing gas in the presence of nitric acid added in an amount within the range from about 0.1 to about 1 gmole per kg of water accompanying the ecllulose pulp at a temperature within the range from about 40 to about 120°C, for an activating time at an activating temperature of from about 15 to about 180 minutes, at an activating temperature of troin 50 to 90°C of from about 5 to about 120 minutes, and at an activating temperature above 90°C of from 1 to about 10 minutes; weating the treated pulp and then subjecting the washed pulp to at least one delignifying stage in an agueous alkaline medium.
 - A process according to claim 1 in which the amount of added mitric acid is within the range from about 0.15 to about 0.80 gmole per by of water.
 - 3. A process according to chim 1 in which the amount of added nitric acid is within the rango from about 0.25 to about 9.60 gmole per kg of water.
 - 4. A process according to claim I in which the temperature is within the range from about 50 to about 100°C.

- 5. A process according to claim 1 in which the temperature is within the range from about 55 to about 90°C.
- 6. A process according to claim 1 in which the delignifying thage is carried out in the presence of coygon gas.
- 7. A process according to ciaim 1 in which the delignifying stage to carried out in the absence of excess gas.
- 8. A process according to claim 1 in which the delignifying stage is carried out in the presence of peroxide bloaching agent.
- 9. A process according to claim 1 in which the nitric acid is added to the celluluse pulp prior to addition of altragen oxide.
- 10. A process according to claim 1 in which the amount of nitrogen exide charged, calculated as monomer, is within the range from about 0.1 to about 4 kilomoics per 100 kg of lighth in the cellulose pulp.
- 11. A process according to claim 1 in which the amount of oxygen added to the pulp in the activating stage is within the range from about 0.08 to about 5 moles O_8 per mole of NO_2 .
- 12. A process according to claim 11 in which the amount of oxygen is within the range from about 0.65 to about 5 males O_8 per mole of NO_2 .
- 13. A process according to claim 1 in which the activation is carried out as a continuous activating stage, and at least the major part of the oxygen gas introduced is added to a zone within the rotention time of the advancing pulp within the range from about 70 to about 100% of the total retention time in the activating stage.

- 14. A process according to chain 1 in which the purp consistency is within the range from about 15 to about 80% during at least 50% of the activating time.
- content of the pulp added to the activating stage, the temperature of said stage and the automats of nitric acid, nitrogen exides and experience are so selected that after ball the activating time has passed the concentration of NO plus NO₂ in the gas plante is not less than 0.05 namely, per liter.
- 16. A process according to claim 1 in which the cellulose pulp is couled to below 40°C during a final part of the activating stage.
- 17. A process according to claim 1 in which the cellulose pulp is flushed from the activating stage with water or an aqueous solution.
- 18. A process according to claim 1 in which the nitric sold to recovered from the pulp leaving the activating stage.

SPECIFICATION

It is well known that chlorine containing bleaching agents give rise to chlorinated aromatic substances and bioaccumulatable chlorinated substances during the bleaching. If these are discharged with waste bleaching liquor into streams and lakes, they are taken up by fish. These substances caunet be destroyed by biological partification of the sewage water. Some chlorinated byproduct substances have been found to be mutagees.

Consequently, disposal of chlorine-containing waste

10 bleaching liquor from bleaching plants constitutes a very serious
problem. Efforts have been made to reduce the use of free or
elementary chlorine by the bleaching of callulose pulp by use of
chlorine dioxide instead. The production of chlorine dioxide
requires about three times as much electrical energy per kilogram

15 of active chlorine as elementary chlorine.

Nitrogen dioxide has been proposed as a substitute for chlorine in the bleaching delignification of cellulose pulp, and has been studied by Clarke (Paper Trade Journal, Tappi. Sect. 118 62 (1944)). Clarke has found that collulose pulp can be partially delignified by treating the pulp in an aqueous suspension for from 1 to 1.5 hours at 90°C with nitrogen dioxide, followed by extraction at 90°C for 30 minutes, or at 50°C for 60 minutes at a 7% pulp consistency and an alkali charge corresponding to 2% NaOil, calculated on the day weight of the pulp. The treatment results in

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a severe depolymerization of the cellulose, which is reflected in a very low viscosity of the treated pulp, compared with pulp subjected to chlorination and alkali extraction.

Bourtt (French patent specification No. 2, 158, 873) avoids depolymerization by applying a delignification process in which the pulp is treated with nitrogen diaxide at low temperature, proferably a temperature below 20°C, and for a long partod of time, followed by an alkali extraction under mild conditions. The cellulose pulp is only delignified to a very small extent, however, and the method does not alford any solution to existing environmental problems.

The delignification of lignocellulosic material by treatment with nitrogen dioxide, followed by washing with water, treatment with electi, and subsequent treatment with oxygen gas, has also been proposed in Swedish patent application No. 77 05138-5. However, this technique has not been put into commercial practice, because although enabling a high dogree of delignification, the method causes a drastic lowering of the viscosity.

Another proposal which has not come into practice has been made in Swedish patent application No. 75 06846-4. This bleaching process includes the steps of (1) treating the calmiosic material with a blend of nitrogen monoxide and nitrogen dioxide with nitrogen monoxide in a molar excess, (2) washing with water, and (3) then treating with alkali, for example, in the presence of exygen gas, under superatmospheric pressure. The nitrogen dioxide can

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uptionally be generated in situ from nitrogen monoxide and oxygen, in which case the nitrogen monoxide is added in an excess of four times the added molar amount of exygen. The reaction proceeds under superatmospheric pressure with respect to nitrogen monoxide; for example, 7 kp/cm² is shown in Example 1. The nitrogen exides are removed by depressurizing, followed by evacuation. In every Example, a superatmospheric pressure is employed in the handling of the nitrogen exides. The handling problems remain, with a great risk of injury to both the internal and external surroundings, and a 10 high consumption of nitrogen exides. This medical abso results in a considerable lowering of the viscosity, although it does enable a high degree of delignification to be obtained.

When the pretreatment with nitrogon oxide is followed by an oxygen gas bloaching stage, it is said to be suitable, subsequent to displacing or washing from the pulp pulping liquor derived from a pulping process with the use of weste liquor derived from the oxygen gas bleaching, to wash the pulp with the acid washing liquid obtained in the washing stage after the pretroatment. When the acid washing liquid is not washed from the pulp before treating the pulp with 20 mitrogen dioxide, the pH of the liquid is reported to be 2.0, which corresponds to about 0.01 gmole nitric acid, calculated per kg of water in the pulp. The prime object of the method is to remove barmful metal companies from the pulp.

In summary, the pretreatment of celluloso pulp with nitrogen 25 dioxide NO, before an oxygon gas bleaching makes possible a more

complete delignification and an improved oxygen gas bleaching, without deterioration in the paper-making proporties of the pulp.

The patent literature indicates that nitrogen monoxide NO gives a like effect. Mowever, relatively large quantities of nitrogen oxides and starting material (ammonia) for the mambacture of said oxides, respectively, are consumed in the process.

In accordance with Samuelson, Canadian Script No. 379, 102, filed June 5, 1081, the activation stage is carried out with nitrogen dioxide gas in the presence of oxygen gas in an amount such that nitrogen monoxide formed as an intermediate is consumed, while regulating the amount of oxygen gas that is added in such a mannor that at the conclusion of the activation stage practically all of the nitrogen monoxide and altrogen dioxide have been consumed.

That invention accordingly provides a process for the treatment of cellulose pulp with nitrogen dioxide NO₂ adapted for application before an oxygen gas bleaching to make possible a more complete delignification and an improved oxygen gas bleaching, without deterioration in the paper-making properties of the pulp, which comprises subjecting the cellulose pulp to an activation reaction with altrogen dioxide gas in the presence of water and pure exygen gas in an amount within the range from about 0. I to about 5 moles per mole of NO₃ and in an amount within the range from about 0. It to about 5 moles about 5 moles per mole of NO₃, so that altrogen monoxide formed in the activation is utilized in the activation reaction.

That invention also provides a process for the delignification of cellulose pulp, including chemical pulp prepared from the digration of lignocellulosic material, which comprises bringing the collulose pulp in an activation stage in the presence of water and in content with a gas phase containing nitrogen dioxide and modifying the lighth of the cellulose pulp by reaction with nitrogen dioxide; adding exygen gas to the activating reaction in an amount within the range from about 0.1 to about 5 moles per mole of M2 and in an amount within the range from about 0.5 to about 5 moles per mole of NO, so that nitrogen monoxide formed in the activation is afflixed in the activation reaction; and then in a second stage, subjecting the pulp to an exygen gas bleaching in the presence of an alkaline-resulting neutralization medium or neutralizing agent.

In accordance with Samuelson, Canadian patent 1,159,203

15 insuch incomber 27, 1983, cellulose pulp produced by chemically
pulping lignocellulosic material is contacted in an activating stage
in the presence of water with a gas phase containing NO₂ and oxygen
gas, which is supplied in order to utilize the intermediate product
NO for activation; and thereafter the pulp is subject to an alkali
20 treatment, both the activating stage and the alkali treatment stage
being carried out under drastic conditions, at such high temperature
during the activating stage as to obtain a certain degree of degradation of the cellulose molecule, and at a temperature during the
alkali treatment process within the range from about 95 to about 150°C,

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suitably from 101 to 140°C, preferably from 110 to 120°C, the treatment time at 95°C exceeding 45 minutes, at 101°C exceeding 30 minutes, and at 110°C exceeding 15 minutes.

The change in the intrinsic viscosity of the cellulose pole is used as a measurement of the extent to which the cellulose molecules have been degraded. The viscosity values given therein have all been determined without removing lignin and hemicellulose, which is the most reproducible method for pulps with a moderate lighth content (for example with solfate pulps having a Kappa number below 35).

This process has however the disadvantage that it requires a very high alkali charge, and results in a high loss of carbohydrates, if the two-stage process is carried for enough to achieve a low lightn content.

In accordance with Samuelson, Canadian Sorial No. 399, 990,

15 filed March 31, 1982 , residual lights in collulose pulp produced
by chemically pulping lighocollulosic material is removed while
maintaining good pulp quality by contacting the collulose pulp in an
activating stage in the presence of water with a gas phase containing NO₂ and exygen gas at a temperature within the range from about

20 do to about 100°C sufficient to obtain a degradation of the collulose
inclosules resulting in a reduction in the intrinsic viscosity of the
collulose pulp during the activation stage within the range from about 2
to about 35% compared to the intrinsic viscosity prior to the activation;
and then subjecting the pulp to an exygen gas-alkelt-treatment at a

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oxygen partial pressure within the range from about 0.005 to about 0.18 MPm.

A partial pressure of 0.18 MPa with respect to oxygen gas during a major part of the oxygen gas-alkali-treatment affords

5 rapid delignification and good selectivity, while delignification at 0.005 MPa takes place very slowly. At a partial pressure below 0.005 MPa, the bleaching is reduced, and the brightness of the pulp is impaired. The pulp acquires a greyish color, but pulp treated at a higher oxygen gas pressure becomes a pure yellow color. In addition, the pulp yield decreases at low oxygen pressure.

In accordance with the invention, a process is provided for activating chemical cellulose pulp and then delignifying bleaching the activated pulp, which comprises treating chemical cellulose pulp. in an activating stage with altroyer exides in the form of NO, and/or 15 NO and/or polymer forms and double-moleculos thereof, such as N_2O_4 and N_4O_8 , and with an exygen-containing gas in the presence of nitric acid added in an amount within the range from about 0.1 to about 1.0, suitably from 0.15 to 0.80, and preferably from 0.25 to 0.60, gmole per by of water accompanying the collulose pulp at 20 a temperature within the range from about 40 to about 120°C, suitably from 50 to 100°C, proferably from 55 to 90°C for an activating time at an activating temperature of from 40 to 50°C of from about 15 to about 180 minutes, at a temperature of from 50 to 90°C of from about 5 to about 120 minutes, and at tempera-25 turns above 90°C from 1 to about 10 minutes, followed by a washing and at loast one delignifying stage in an aqueous alkaline

medium, either in the presence or in the absence of waygen gas and/or percuide.

The combination of these uitrogen oxides and nitric acid provides an activating effect which results in a greatly improved delignification after the alkaline delignifying stage. The delignifying effect obtained in accordance with the invention with 2% NO₂ by weight of the day pulp is approximately the same as that obtained with twice the amount of NO₂, if no nitric acid is added. This is surprising, since treatment of the pulp with nitric acid in a concentration within the stated range prior to the alkaline stage, without any addition of NO₂ and/or NO, has no appreciable effect on the delignification. The activating effect is obtained irrespective of whether or not expense or peroxide is present in the alkaline delignifying stage.

is present during the activating stage, depolymerization of the carbohydrates, primarily in collabose, is slowed down in the alkaline delignifying stage when the alkaline medium constitutes as exygen gas delignifying medium. Thus, under optimum conditions

20 white there is a certain depolymerization (loss in viscosity) in the activating stage, a pulp is constitutes obtained whose viscosity after the alkalius exygen gas delignifying stage, not only when compared at the same lights content (Kappa number) of the pulp but also when compared at the same reaction time in the exygen gas stage, is

markedly higher than that of paip similarly activated, but without nitric acid being added during the activating stage. Obviously, when an optimum amount of nitric acid is present, the activation provides a chemical roaction which greatly inhibits the degradation of cellulose in the subsequent alkalino oxygen gas bleaching stage.

As a measurement of the degradation of collulose molecules, there is used here the change in the intrinsic ylacosity of the cellulose pulp, determined in accordance with SCAN-C15:62. In the case of fully bloached paper pulps, the viscosity corrustly should not be less than 900 dm^5/kg . All of the viscosities given below have been determined without lighth and hemicellulose having been removed, which is the most reproducible method, in the case of pulps having moderate lighth contents. However, it should be observed that lignin and hemiceliulose contribute only slightly to the viscosity in comparison with the same amount of weight of collulose molecules, and that delignification and bleaching are intended to decrease the light content. An appreciable loss of handcellulose can also occur, while the loss of pure collulose is very small under the conditions used. Consequently, in those cases where depolymerization of the cellulose is negligible, the intrinsic viscosity will increase. With plac suffice palps of the kind used in the majority of Examples given herein, a decrease in the Kappa number of 10 units results in an increase in viscosity of about 50 dm3/kg, under those conditions where depolymerization of the cellulose can be ignored, while a corresponding increase in

the case of sulfite gulp and hardwood sulfate pulp is markedly higher because more hemicellulose is lost.

The nitrogen dioxide can be supplied as substantially pure NO_2 . NO_3 can also be formed in sim in the reactor upon charging attric oxide and expenditure. NO_3 plus NO may also be supplied. Dinitrogen televaide (N_1O_3) and other polymeric nitrogen dioxide forms are considered to be included in the term "nitrogen dioxide" (NO_2) . One make of dinitrogen tetroxide is considered to be the same as two moles of nitrogen dioxide. Adducts in which nitric oxide is present are considered in the same manner as nitric oxide. Thus, dinitrogen trioxide (N_2O_3) is considered as one mole of nitric oxide and one mole of nitrogen dioxide. Adducts in which expenditure the present probably occur as intermediates.

The amount of nitrogen oxides charged is adapted to the lightn content, the desired degree of delignification, and the tolerable attack on the carbohydrates. Calculated as monomers, the amount is normally within the range from about 0.1 to about 4, satisfully from 0.3 to 2, preferably from 0.5 to 1.2, kilogrammoles calculated per 100 kg lightn in the pulp entering the activating stage.

A certain amount of oxygen gas must be supplied to the activating stage, both when nitrogen dioxide (NO₂) is charged and when nitric oxide (NO) is charged. This oxygen-containing gas may be air.

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In order to obtain the best possible results with the simplest apparatus possible, it is suitable to supply the exygen to the activating singe in the form of a substantially pure exygen gas.

Liquid exygen can also be supplied to the activating singe, and vaporized, e.g., when entering the reactor in which the activating process is carried out. The use of substantially pure exygen results in a lower content of NO plus NO₂ in the gas phase than when air is used. It also means that only a minor quantity of heart gas needs to be removed from the reactor, and optionally treated to reader to residual gases harmless.

The amount of oxygen added to the activating stage is solveted according to the amount of nitrogen exides added, so that the addition of O_r per mole of NO₂ amounts to si least 0.08, suitably from 0.1 to 2.0, preferably from 0.15 to 0.30, mole.

15 If NO or a mixture of NO and NO₂ is used instead, the expen gas addition is so made that the amount of exygen is at least 0.60, suitably from 0.65 to 3.0, preferably from 0.70 to 0.85, mole O₂ per mole of NO charged. When NO is used the charge is preferably made batchwise or continuously in a manner such that exygen is complied in portions or continuously before the supply of NO is terminated. In this way, activation is more uniform than when exygen gas is not supplied until all NO has been charged to the reactor. The reactor can be designed for batchwise operation, or for continuous operation in a reaction zone with continuous food-in,

transit through and delivery of cellulose pulp from the continuous zone, and the supply of gases thereto.

Nitric acid can be added to the entiruting stage when activation with nitrogen oxider and with oxygon-containing gas has been completed. For example, the cellulose pulp can be flushed out from the activating reactor vessel or from a zone thereof with nitric acid. Nitric acid can also be supplied during treatment with the aforementioned guses.

It has been found most suitable for delignification to supply

the acid to the pulp before the pulp is brought into contact with

nitrogen oxides. Impregnation with excess nitric acid and romoval

of the excess, for example, by filtration and/or pressing, is a

proformed embodiment. Irrespective of how the acid is added, it
is advantageous for the pulp containing the nitric acid to be subjected

to activation with nitrogen exides and exygen-cuntaining gas at a pulp

consistency within the range from about 15 to about 80%, suitably

from 20 to 90%, proferably from 25 to 45%. The stated pulp

consistency should be maintained for at least 50% of the colivating

time. The pulp consistency can be malatained over the whole of

the activating time to great advantage. When the supply of gases to

the activating stage has been substantially completed, dilution with

water or preferably nitric acid can be made, which can afford

advantages in the case of certain types of pulp.

One characteristic result of the process according to the invention is that more eitric exide and nitrogen dioxide is obtained in the gas phase dish when no altric acid is present, with the same addition of altrogen exides and exygen gas, and the reaction parameters in other respects also being equal. An increased pulp consistency and an elevated temperature also result in an increase in residual NO_x and/or NO gas content. Tests have shown that the moisture content of the pulp, the temperature of the activating single and the addition of nitric acid, altrogen exides and exygen gas should be so adapted that effor one-half the activating time, the amount of the place NO_x in the gas phase amounts to at least 0.05 manuals per producing pulps having a high degree of delignification, the amount of NO plus NO_x in the gas phase should be at least 0.1 and preferably at least 0.15 manuals per liter.

It has been found that the major part of the nitrogen axides added are consumed very rapidly when there is a surplus of oxygen gas in the reactor vessel, and that the rate of consumption is very slow towards the end of the activation period. This is due to the fact that aftric oxide is split off from the celiulose pulp as a result of some unknown reaction. This previously unknown reaction is favored in some way by the nitric acid present, and seems to explain the surprising technical effects achieved by the addition of nitric acid in accordance with the invention. The presence of exygen gas by a

preroquisite for achieving these effects.

It is particularly suitable to charge NO₂ and/or NO in the vicinity of the feed-in or inlet end of a continuous activating stage. In order to obtain the best possible activation and utilization of the altrogen exides amplied, and the least possible gas efficient, and as little treable as possible with rendering unconsumed nitric exide and nitrogen dioxide harmless, in the case of a continuous activating stage, expenses, preferably the major part of the expenses, should be fed in at one or more zones located in the vicinity of the outlet end of the reactor. Satisably, the expenses is supplied to a zone which is so located that the retention time of the advancing pulp corresponds to 70 to 100, anitably 80 to 100, preferably 90 to 160%, of the total retention time in the activating stage.

It has also been found advantageous to lower the temperature of the collulose pulp later in the activating stage, for example, when 80% of the activating time has passed. This lowering of the temperature may advantageously be so effected that the temperature of the pulp is less than 40°C, for example, within the range from about 10 to about 35°C, suitably from 20 to 30°C, and the releation time at a temperature below 40°C is, for example, from 10 to 120 minutes, preferably from 15 to 60 minutes.

The time at which the temperature is bolow 40°C is not included in the time ranges above. Any cooling required to reduce the temperature can be effected indirectly, for example, by cooling the gas phase, or by introducing cold oxygen, for example, liquid exygen, to the activating stage. Water can also be evaporated by lowering the pressure.

When activation is effected continuously, the feed-in of inert gas into the reactor and the discharge of gas from said reactor should effectively be prevented. This can be achieved by providing 10 known gas slutens for the pulp at the feed-in and and at the feed-out and of the vessel.

The total gas pressure (including steam) in the reactor is suitably held at about atmospheric pressure, preforably at a low subatmospheric pressure corresponding, for example, to from 15 about 0.001 to about 0.01 MPa.

It has been found particularly suitable to discharge the cellulose polp from the activating stage by flushing with water and/or an aqueous solution. In accordance with one preferred embeddment, the flushing water is waste liquor recovered from the process and 20 containing nitric acid and an organic substance.

The major part of the nitrogen exides supplied in accordance with the method according to the invention gives rise to altric anid.

According to a presented embodiment, the affric acid used in the solivating stage is recovered either completely or partially from the pulp discharged from the activating stage. The aitric acid can be recovered in a known manner, for example, by washing and/or displacement. The acid can also be recovered by pressing the pulp, professily subsequent to dilution with water and/or an aqueous solution. Adventageously, recovery of the acid is offected in accordance with the countercurrent flow principle, such that, after the activating stage, the pulp is brought into contact with waste

According to a further embodiment, which has been found to provide a better result than when puro nitric acid is charged to the activating stage, the wasto liquor containing after acid recovered from the activating stage is used to improposts the collulose pulp added to the activating stage. Imprognation of the pulp is suitably effected by passing the ingoing pulp in countercurrent flow with the wasto liquor, so that the pulp is progressively brought into contact with waste liquor of increasing nitric acid concentration. This countercurrent imprognation of the collulose pulp is preferably carried out when substantially all cooking liquor originating from the cooking of the cellulosic raw materials has been washed or displaced from the pulp.

In accordance with a preferred embediment, the cooking liquor present in the pulp is washed or displaced therefrom with

waste liquor obtained from the alkaline delignifying stage, this waste liquor being substantially removed when the pulp is impregnated with waste liquor from the nativaling singu-

When ammonia is need to produce altric oxide and/or nitrogen dioxide, aftric acid is formed as a hyproduct. This aftric acid can be used to advantage in the activating process, preferably in combination with waste liquor recovered from the activating stage. The preparation of mitric oxide and nitric acid in situ enables a very uniform reaction to be achieved during the activating process, so which can readily be governed through a controlled successive supply of nitric oxide and oxygen, so that the whole of the cellulose pulp comes into effective contact with the gases, and so that local overheating within the reactor is avoided.

By the process according to the invection, it is possible to remove at least 75% of the lighth remaining in the pulp after the pulping, while maintaining good pulp properties. The residual lighth is removed by treating the pulp in accordance with known final bleaching techniques, for example, by using as bleaching agents chlorine dloxide, hypochlorite and optionally, chlorine.

A number of advantages are obtained when treating cellulose pulp in accordance with the invention. The most important of these advantages is that the cost of chemicals for the pollunting stage is greatly reduced, in comparison with previously known techniques. This saving in chemicals is expressed in different ways, depending on whether commercial nitrogen exides are used (nitrogen

dioxide can be bought as a commorpial product) or whether the nitrogen exists are produced from ammonia by oneself.

Thus, when practicing the process according to the invention, and using commercial nitrogen dioxido, the amount required is only balf that which would otherwise be necessary. The nitric acid required is obtained more or less free, since nitric acid is generated during the activating stage, and can be recovered upon the completion of said stage. Initially, it may be necessary to purchase fresh nitric soid, but the amount required will never be cont-significant.

If the nitric oxide and/or nitrogen dioxide is produced in-house, using ammonia as a starting chemical, a considerable amount of nitric acid is formed at the same time. The invention makes use of this nitric acid, which would otherwise be difficult to dispose of. Thus, the installation of a plant for the production of nitrogen axides in connection with the blooch plant is a feasible proposition, and reduces the costs for activating chemicals to a minimum.

The following Examples represent preferred embediments 20 of the invention.

EXAMPLES 1 to 8

Activating Stage:

A pine sulfate pulp having a Kappa number of 33.5 and an intrinsic viscosity of 1105 dm³/ky was pressed to a solids content of 39%. The pulp was then impregnated with nitric acid by mixing for ten minutes at room temperature with an amount of nitric acid corresponding to 0.4 g mole per ky of water in the pulp. As a rosult, the consistency of the pulp was reduced to 30%.

The pulp was charged to a rotary reactor, which was then

10 evacuated and heated to a temperature of 58°C over ten minutes.

2% NO₈ by weight of the bone dry pulp was then charged by vapor
tzing liquid N₂O₄ in the evacuated reactor. Oxygen gas was charged

to the reactor in three portions over a period of two minutes to

bring the pressure in the reactor to atmospheric prossure. Five

minutes after the charge of NO₈ was begun, the temperature was

reduced to 50°C, and this temperature was maintained for 55 minutes.

The reactor was then cooled to 30°C over lifteen minutes, and then

the pulp was finished from the reactor with cold water, halting the

treatment. The total time for the treatment was thus 75 minutes,

20 including the time taken to cool the reactor. These are Examples

1, 2, 3 and 4.

Four additional experiments were made in accordance with the invention, and these are Examples 5, 6, 7 and 8. These activating treatments were carried out in a similar manner to

Examples 1 to 4, except that the temperature was maintained at 50°C over the whole of the activating stage.

Four Control activating treatments were made, these being denoted A, B, C and D, under conditions the same as those in Examples 5 to 8, except that water was added to the pulp prior to the activating stage instead of nitric acid.

Four Control activating treatments were also made,

Controls F., F. G and H., with a charge of 4% NO₂ without the addition
of nitric acid. The same conditions after the activating stage were

10 used as those in the others. The whole NO₂ charge was then made

over a period of two minutes, and then exygen gas was charged in

three portions for two minutes, to bring the pressure to atmospheric

pressure. The temperature was kept constant at 50°C for 75 minutes.

Four additional Control activating treatments, Controls I, I,

15 K and L were made with a first charge of 2% NO₂ at 55°C. After five
minutes the pulp was cooled to 50°C and a further 2% charge of NO₂

was made. After fifteen minutes, calculated from the time at which
the charge of NO₂ was begun, O₂ was charged in three portions for
two minutes, to bring the presence to atmospheric pressure. After

20 a total activating time of sixty minutes, the poly was cooled to 30°C.

After a total time of 75 minutes, the activating process was interrapted as before.

Table I shows the most important parameters in the activating stage, and the pulp properties achieved thring Brese

experiments, all of which were catried out at a total time of 75 minutes.

In order to estimate the degradation of the enzhohydrates and the formation of readily soluble light during the activating stage, samples of the pulp were taken after the treatment, and washed with 0.2 M aqueous NaIICO₃ solution at room temperature, and then with water. Subsequently, the samples were dried rapidly in a stream of air at 35°C. The results of these are given in Table I.

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	After NaHCO, wash	Viscosity dm3/kg	1200	1038		1161	1124	1137
	After Nai	Keppe. Number	22.3	32.7		27.9	23.0	25.0
н.	NOcontent &	in the cas chese amole/i	0. 12	0, 19		0.02	p. 18	0.09
TABLE 1	Hohest	temperature C	58	. 20		50	Q:	92
	Charged Amount	g mole/	0.4	0.4		0	D	0
i	Chare	og er	ca.	EX.		N	잭	ঝ
		Exemple Mo.	1 to 4	5 to 8	Controis	A to D	可 35 34	7. 0 ² 1
		دی	s				11	•

Measured at the end of the treatment.

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Alkaline oxygen bleaching stage:

The activated pulp from each of Examples I to 4, 5 to 6, and Controls A to D, E to H, and I to L was then weahed with cold water and separated into batches corresponding to the Examples in these five groups. These batches were each subjected to an oxygen gas bleaching proceed at a pulp consistency of 25% for a period of sixty minutes and at a temperature of 106°C. Pure oxygen gas was used, and the oxygen partial pressure was 0.11 MFs, measured at 106°C. Alkali in the form of fore sodium hydroxide was charged to the batches in each group in a quantity of 1.0, 1.5, 2.5 and 4.0%, respectively, as shown in Table II, by weight of the dry unbleached pulp. In addition, there was charged to each batch magnesium complex with spont bleaching liquor in an amount corresponding to 0.2% Mg, calculated in the same manner as above.

TABLE II

	Oxygen gas bleached pulp produced with varying charges of NaOII		
Example No.	NaOH 栄_	Kappa Number	Viscosity dm³/kg
1 .	1.0	13.2	1038
2	1.5	10.8	1031
3	2.5	0.1	1014
4 ·	4.0	8.2	936
5	1.0	13.9	1034
6.	1.6	11.7	1040
7	2.5	9.3	1011
8	4.0	8.5	941
Controls		•	
A	1.0	31. 0	1078
В	1.5	15.0	1088
C	2.5	12,4	990
D	4.0	10.4	845
E .	1.0	15.8	1045
F	1.5	12.7	1.055
G	2.5	10.4	1034
H	4.0	R.B	953
1	1.0	10.1	1050
J	1.5	12.9	1057
K	2.5	10.R·	1038
L	4.0	9:8	935

the pulp was washed with sodium higarhonale, an insignificant reduction in the intrinsic viscosity was obtained when mitric acid was not added prior to the NO₂/O₂ treatment process. Furthermore, a lower content of NO₂ in the gas phase was obtained than in Examples 1 to 8. In Examples 1 to 8 the viscosity was markedly lowered, which is of course a disadvantage, as is also the higher residual NO₈ gas content. An increase in the formation of readily soluble lights is reflected in the lower Kappa number of the pulps to washed with sodium bicarbonate, which pulps were activated in the presence of other acid, than in Controls A to D in the presence of water.

delignification, in comparison with Controls A to D after the oxygen
15 gas stage, particularly in the case of the lower charges of NaOct.
These pulps were impregnated with nitric acid prior to charging
altrogen dioxide. The effect was so great that the selectivity,
defined as the viscosity at a given Kappa number, was much higher
in Examples 1 to 8 than in the Controls A to L, in which so nitric
20 acid was used. It is of particular interest to note that when the
larger quantities of sodium hydroxide were charged (2.5 and 4%,
Examples 3, 4, 7 and 8) the viscosity of the oxygen gas blenched
pulps, despite the marked lowering of the oxygen gas blenched
cativating stage, when said stage was carried out after adding nitric

soid, was appreciably higher after a given period of time with constant conditions in the exagen gas bleaching stage.

The experiments show firstly that in accordance with the invention there is obtained an improved designification, and secondly that the special conditions during the activating stage result in a strong returdation of the depolymentation of the carbohydrates, primarily depolymentation of the cellulose, during a subsequent oxygen gas bleaching stage.

A somewhat higher selectivity and lower Kappa number with
the same alkali charge was obtained according to the invention, when
the highest temperature was 58°C and the temperature lowered
during the activating stage (Examples 1 to 4) than when the entire
activating stage was carried out at a temperature of 50°C
(Examples 5 to 6). In addition, far loss nitrogen dioxide was obtained
in the gas phase at the end of the activating time, which represents
an approxiable advantage with respect to pollution of the environment.
Separate experiments have shown that a further reduction in the
temperature to 22°C and an increased contact time between the gas
phase and the pulp at a low temperature in the final stage of the
20 activating stage (after treatment) results in a further lowering of the
residual gas content.

As expected, in Controls E to 1, with 4% NO, a much higher degree of dollgoiffention was obtained than when using 2% NO.

The Kappa numbers, however, were consistently higher than those

reached after the same treatment process in the alkali slage, when the protreatment process was carried out in accordance with the invention with half the amount of NO, charged. Compared at the same Ruppa number, the differences in viscosity between the Controls using 4% NO, and Examples 1 to 3 using 2% NO, were hardly sigafficant. Since nitric acid is formed by the attrogen diexide charged, and can be recovered for use in the process, the invention enables a reduction by about 50% in the charge of chemicals to the activating stage, in comparison with previously known techniques.

In another series of Controls, Controls M, N, O and P, the pulp was impregnated with hydrochloric acld lastesd of nitric acid, to give an HCl concentration of 0.4 g male per by of water in the impregnated pulp. In other respects, the Controls were carried out in the same manner as the Examples 5 to 8. After washing with 15 sodium bicarbonate, the Kuppa number was only 1.2 units lower than in the Controls A to L carried out with water, i.e., markedly higher than in Examples 1 to 8, using nitric and, while the viscosity was 22 units lower than in corresponding Examples 1 to 8 with nitric acid.

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After oxygen gas bleaching the paip, the Kappa number 30 differed on average by less than 5½ from the Centrals A to L with water, a difference which has hardly any significance, while powerful degradation of celluloso was reflected in the viscosity values, which were 66 to 86 units lower than in Controls A to L using water. Thus, the selectivity was even worse than Controls

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A to L., with no acid present. Consequently, nitric acid cannot be replaced with hydrochioric acid or any other acid. The Controls show that the more effective delignification obtained in the presence of nitric acid at the given concentrations is not dependent on an acid hydrolysis of lighth bonds.

EXAMPLES 9 to 14

A satisfic pulp of softwood, mainly pine, having a Kappa number of 30.7 and an intrinsic viscosity of 1225 dm²/kg, was present to a solids content of 31%. The mass was then treated in a glass reactor with 4% NO₂ by weight of the dry pulp at 57°C. The reactor was avacuated and heated to 57°C prior to charging vilrogen dioxide by vaporizing liquid N₂O₄. The charge of NO₂ required three minutes. Oxygen gas was charged to the reactor in small portions over a period of two minutes, to bring the reactor pressure to 10 atmospheric, and the reactor was related for a total reaction time of fifteen minutes.

The activating process was continued by diluting the pulp (without previous washing) with nitric acid of varying conventrations, so that an 8% solids content was obtained. The pulp was permitted to react with the charged nitric acid at varying temperatures and times. The nitric acid was then filtered off, and the pulp washed with cold water.

The pulp was then subjected to a delignifying stage, using a not alkali treatment in the absonce of air or oxygen gas at a pulp 20 consistency of 24%. The alkali charge comprised 5% NaOII, by weight of the bone dry pulp. The temperature was 105°C, and the time 45 minutes. After the hot atkali treatment the pulp was washed with water, and analyzed.

In addition, three Controls, Control Q. Control R and 25 Control S, were run. In Controls Q and R, the pulp subsequent to being

treated with NO₂/O₂ was diluted with water to a consistency of 8% and the suspension maintained at 40°C and 60°C, respectively, for starty minutes.

In Control S, the pulp was washed with cold water and subjected to a hot alkali treatment process immediately after being treated with NO₂/O₃ for fifteen minutes at a consistency of \$1%.

Controls Q and H showed no appreciable change in Kapps number compared to Control S, although a slight lowering of the viscosity compared to Control S can be seen.

10 The differing parameters during the activating stage and the pulp properties obtained are set forth in Table III.

TABLE III

					Analysia of hot aikali treated pul		
5	Example No.	Charged HNO.		Time Minutes	Kappa Number	Viecusity (dm³/kg)	
	9	Q. 2	40	60	12.2	1165	
	1.0 .	0.2	60	60	9.1	1127	
	11 .	0.4	. 40	60	10.0	1200	
	12	0.4	. 60	30	10.0	1166	
10	13	0.4	ថម	80	7.8	1094	
	. 1.4	6, ∳	60	120	6.8	1946.	
	Control Q	ņ	40 .	60	12.7	1208	
:	Control R	o .	GO	. 60	12.4	1176	
	Control S			- ' .	12.5	1214	

In Examples 10 to 14 in accordance with the invention, a significant lowering of the Kappa number was obtained compared to Control 6. The effect was much greater at 60°C (Examples 10 and 12 to 14) then at 40°C (Examples 9 and 11). At 60°C there was obtained a marked effect with 0.2 M attric soid, while the effect was small at 40°C at this concentration. An increase in the concentration and in the treatment time resulted in improved delignification, but at the same time in a decrease in the viscosity, which, however, was nuclerate in comparison with the decrease in the Kappa number.

Normally, a viscosity of 900 dis /kg is considered a minimum for a biesched paper pulp of high quality. When the concentration of altric acid was higher than 1.0 g molo per kg of water, the viscosity decreased below this value under those conditions with respect to time and temperature where a significant effect with 15 respect to delignification was achieved because of the presence of the nitric acid. Thus, when the nitric acid concentration was 1.1 g moles per kg of water, the activating temperature 50°C, and treatment time thirty minutes, the intrinsic viscosity was 840 dms/kg.

Treatment nurried out in accordance with Example 14 results

20 In a significant dissolution of homicallulose, which is advantageous
in the case of special pulps, e.g., in the case of special paper which
is required to be particularly resistant to ageing, while a treatment
time of 120 minutes with nitric acid (0.4 g mole per kg water) at 60°C
is, in general, much too long to obtain an optimum effect.

EXAMPLES 15 to 23

Activating staye;

A sulfate pulp produced from a mixture of 50% sprace,

40% plue and 1.0% aspen, having a Kappa number of 34.8 and an

intrinsic viscosity of 1196 dm²/kg, was pressed to a solide content

of 39%. The pulp was then imprograted with aftric acid by mixing
for ten minutes at room temperature with an amount of nitric acid

corresponding to 0.3 g mode in Examples 15 to 18 and 0.4 g mole in

Examples 19 to 22, per kg at water in the pulp. As a result, the

The pulp was charged to a rotary reactor, which was then exactated and heated to a temporature of 58 °C over ten mirales.

2% NO, by weight of the bone dry pulp was then charged by vaporizing liquid N₂O₂ in the execuated reactor. Oxygen gas was charged to the reactor in three portions ever a period of two minutes to bring the pressure is the reactor to atmospheric pressure. Five minutes after the charge of NO₂ was begun, the temporature was reduced to 55°C, and this temperature was maintained for 30 minutes. Then, the pulp was flushed from the reactor with cold water, halting the treatment.

Four Control activating treatments were made, Controls:

T and U being under the same conditions as above, except that water was added to the pulp pater to the activating stage instead of nitric acid, and Controls V and W being under the same conditions as above.

except that 0.08 g mole nitric acid per kg of water was used.

Four Controls activating treatments were also made, Controls X, Y, Z and ZZ, the first two with a charge of 0.4% nitric acid without the addition of NO, and Controls Z and ZZ without either NO, or nitric acid. The other conditions were the same.

Table IV shows the most important parameters in the activating stage.

		(Tlane		TABLE	<u>tv</u> .
ច្	Example No.	110,	HNO. g mole/ kg H.O	Flighest temperature C	MD ₂ -content in the gas phase mmole/1
•	15 to 18	2	0.3	55	0, 25
	19 to 22	2	0.4	55	. 0.32
	Controls	•	•	•	
	T and U	2	a	55	o. m
10	V and W	2	0.08	55	0.04
	X and Y	0.	0.4	. 55	0.D1
	Z and ZZ	۰ ۵	O	5 5	Ð

Intersured at the end of the treatment

Alkaline cayeen bleaching singe:

The activated pulp from each of Examples 15 to 13, 10 to 22, and Controls T and U, V and W, X and Y, Z and ZZ, was then washed with cold water and separated into batches corresponding to the Examples in these all groups. These batches were each subjected to an exygen gas bleaching process at a pulp consistency of 26% for a period of sixty minutes and at a temperature of 106°C. Pure exygen gas was used, and the exygen partial pressure was 0.11 MPa, measured at 106°C. Alkali in the form of pure sodium.

10 hydroxide was charged to the batches in each group in a quantity of 1.0, 1.5, 2.5 and 4.0%, respectively, as shown in Table V, by weight of the dry unbleached pulp. In addition, there was charged to each batch coagnesium complex with spent bleaching liquor in an amount corresponding to 0.2% Mg, calculated in the same manner as above.

TABL); V

		oten head designed pulp produced with verying HOMM to engree			
5	Example No.	МаОн Ж	-Kappa Mumber	Viscosily dm:/kg	
	1 5 .	1.0	16.8	1126	
	18	1.5	13.7	1009	
	17	. 2.5	11.3	1061.	
10	18 .	4,0	10, 1	884	
•	19	1.0	14.5	1069	
	2.0	1.5	12.0	1042	
•	. 21	2.5	10.0	994	
	22	4.0	8.B	972 :	
1.5	Controls			:	
	${f T}$	1.5	17.4	1040	
	Ů.	4.0	12.2	928	
	Υ	1.5	17.5	1047	
	W	4.0	12.0	920	
20	· x	1.5	22.3	908	
	У·	4.0	14.1	992	
	7	1.5	22.1	991	
	2.7.	4. 0	14.4	931.	

As is shown in Table V, delignification was much poorer in Controls T, U, V and W compared with Examples 15 to 22 in accordance with the invention. In the Controls, the concentration of nitrogen dioxide in the gas phase was much lower than in the banmples. Like the gas analysis given in Table I, these analyses indicate that the nitric acid charged in the Examples carried out in accordance with the invention contributes in an enknown manner to increasing the content of components in the gas phase which promote the activaling process.

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10 Controls X and Y show that the introduction of nitric acid without the addition of nitrogen diexide has an insignificant effect on delignification, under the conditions used in the Examples in accordance with the invention. The results differ insignificantly from those obtained in Controls Z and ZN, where the subside pulp was expenses gas bleached directly, without any preceding activation.